10/772,235 UPDATED, EXPANDED STN SEARCH TRANSCRIPT

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CA/CAplus Indian patent publication number format defined RDISCLOSURE on STN Easy enhanced with new search and display
                                                                                                               RDISCLOSURE reloaded with enhancements
JICST-PENUS removed from database clusters and STN.
GENBANK reloaded and enhanced with Genome Project ID field
CHEMCATS enhanced with 1.2 million new records
CA/CAPLUS enhanced with 1870-1889 U.S. patent records
INPADOC replaced by INPADOCDB on STN
New CAS web site launched
                                                                                                                                                                                                                                                                                                                                                                           TOXCENTER enhanced with BIOSIS reload CA/CAplus enhanced with additional kind codes for German
                                                                                                                                                                                                                                                                                                                                                                                                                                             CA/CAplus enhanced with IPC reclassification in Japanese
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                CHEMCATS accession numbers revised CA/CAplus enhanced with utility model patents from China
Web Page for STN Seminar Schedule - N. America VPIDS/WPIX chanced with new FRAGHITSTR display format CASREACT coverage extended MARPAT now updated daily
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                     CACOMPLUS enhanced with pre-1967 CAS Registry Numbers STN Viewer now available STN Express, Version 8.2, now available LEMBASE coverage updated LAMEDLIRE coverage updated SCISEARCH enhanced with complete author names
                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                                  29 JUNE 2007: CURRENT WINDOMS VERSION IS V8.2, CURRENT MACINTOSH VERSION IS V6.0c(ENG) AND V6.0JC(JP), AND CURRENT DISCOVER FILE IS DATED 4 MAY 2007.
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FILE 'HOME' ENTERED AT 10:08:46 ON 06 JUL 2007

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chain nodes :

ring nodes:

6-19 7-8 7-15 8-9 8-10 9-13 9-19 5-7 ring bonds: 1-2 1-6 2-3 3-4 4-5 4-14 5-6 10-11 11-12 12-13 14-15

exact/norm bonds:
4-14 5-7 6-19 7-8 7-15 9-19 14-15 14-16
normalized bonds:
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containing 1:

G1:0,S

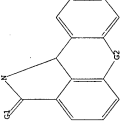
G2:0,S,N

Match level : 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:CLASS 19:Atom

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G1 0,S G2 0,S,N

Structure attributes must be viewed using STN Express query preparation.

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chain nodes :

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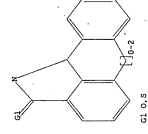
1-2 1-6 2-3 3-4 4-5 10-11 11-12 12-13 isolated ring systems:

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Match level : 1:Atom 2:Atom 3:Atom 4:Atom 5:Atom 6:Atom 7:Atom 8:Atom 9:Atom 10:Atom 11:Atom 12:Atom 13:Atom 14:Atom 15:Atom 16:CLASS 19:CLASS

STRUCTURE UPLOADED

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0 ANSWERS 67 ITERATIONS 100.0% PROCESSED SEARCH TIME: 00.00.01

1831 ONLINE **COMPLETE**
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849 TO 183. FULL FILE PROJECTIONS: PROJECTED ITERATIONS: PROJECTED ANSWERS:

=> S 11 SSS FULL FULL SEARCH INITIATED 10:09:43 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 126 TO ITERATE OR L2 0 SEA SSS SAM L1 F3

5 ANSWERS 126 ITERATIONS 100.0% PROCESSED SEARCH TIME: 00.00.01

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680 TO ITERATE => S L2 SSS FULL FULL SEARCH INITIATED 10:09:48 FILE 'REGISTRY' FULL SCREEN SEARCH COMPLETED - 680 TO ITE

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TOTAL SESSION 343.96 SINCE FILE ENTRY 343.75 => FILE CAPLUS COST IN U.S. DOLLARS FULL ESTIMATED COST

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=> S L4 L6

4 L4

=> D 1-4 IBIB ABS HITSTR

COPYRIGHT 2007 ACS on STN CAPLUS 2004:792352 ANSWER 1 OF 4 CAPLUS ACCESSION NUMBER:

141:424132
Synthesis of chromeno[4,3,2-cd]isoindolin-2-ones DOCUMENT NUMBER:

chromeno[4,3,2-de]isoquinolin-3-ones. Electrophilic versus anionic cyclization of carbamates Carmen de la Fuente, M.; Dominguez, Domingo AUTHOR(S): CORPORATE SOURCE:

Facultad de Quimica, Departamento de Quimica Organica y Unidad Asociada al CSTC, Universidad de Santiago de Compostela, 15782, Spain Terrahedron (2004), 60(44), 10019-10028 CODEN: TETRAB: ISSN: 0040-4020

Elsevier B.V. DOCUMENT IYPE: LANGUAGE: PUBLISHER: SOURCE:

English

OTHER SOURCE(S):

CASREACT 141:424132

AB The total synthesis of chromeno[4,3,2-cd]isoindolin-2-ones and chromeno[4,3,2-de]isoquinolin-3-ones [rom 4-methoxy-9H-wanthen-9-one is reported. The construction of the nitrogenated ring was attempted by both intramol. electrophilic and anionic cyclizations of the corresponding carbamate precursors. Only anionic cyclizations of the corresponding carbamate precursors. Only anionic cyclization was possible for isoindolinones, but for isoquinolinones the electrophilic and anionic routes both afforded excellent yields.

IT 794513-42-1P 794513-43-P 794513-44-3P 794513-44-49

RL: SPN (Synthetic preparation) PREP (Preparation) (preparation of chromeno[4,3,2-cd]isoindolin-2-ones and chromeno[4,3,2-cd]isoindolin-2-one via intramol. electrophilic and anionic cyclization reactions)

Z 23

7945<u>13-</u>42-1 CAPLUS 2H-[1]Benzopyrano[4,3,2-cd]isoindol-2-one, 1,10b-dihydro-5-methoxy-1-methyl- (9CI) (CA INDEX NAME)

RN 794513-43-2 CAPLUS
CN 2H-[1]Benzopyrano[4,3,2-cd]isoindol-2-one, 1,10b-dihydro-10b-d-5-methoxy-1methyl- (9CI) (CA INDEX NAME)

RN 794513-44-3 CAPLUS CN 2H-[1]Benzopyrano[4,3,2-cd]isoindol-2-one, 1,10b-dihydro-5-methoxy-l-methyl-10b-phenyl- (9CI) (CA INDEX NAME)

RN 794513-45-4 CAPIUS CN 2H-[1]Benzopyrano[4,3,2-cd]isoindol-2-one, 10b-ethyl-1,10b-dihydro-5methoxy-1-methyl- (9CI) (CA INDEX NAME)

REFERENCE COUNT: 50 THERE ARE 50 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE RE FORMAT

L6 ANSWER 2 OF 4 CAPLUS COPYRIGHT 2007 ACS on STN ACCESSION NUMBER: 2003:92405 CAPLUS

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PRIORITY APPLN. INFO.:				1997-92254 1998-47502 1998-14518 1998-US181	-922548 -47502 -145181 -US1818	ი	22 4 3		19970903 19980325 19980901 19980902	2223
OTHER SOURCE(S): GI	MARPAT	138:137290								

AB Title compds. [I; Y = alkylhalo, alkyl-COG, COG, bond, CO, O, NRII, CR8; G = NRII R16, OR9, SR9, R10; Z = O, S, NRII; X = NR16, O, S, CR12R13, CO, bond, CR12:CR13, CR12 R13CR14R15; R1-R8, R10, R12-R15 = H, halo, alkylhalo, OH, alkyl, alkenyl, cycloalkenyl, cycloalkenyl, aryl, amino, alkylamino, NO2, nitroso, COZH, aralkyl; R9 = H, OH, alkyl, alkenyl, cycloalkyl, cycloalkyl, aryl, amino, alkylamino, COZH, aralkyl; R11, R16

= H, halo, alkylhalo, OH, alkyl, alkenyl, cycloalkyl, cycloalkenyl, aryl, amino, alkylamino, COZH, aralkyl; the alkyl, alkenyl, cycloalkyl, cycloalkenyl, aryl, aralkyl groups may be substituteds with provises), were prepared Thus, 9-xanthenylmethyl isocyanate (preparation given) was

The latter inhibited PARP in polyphosphoric acid at 90° to give 1,11b-dihydrobenzopyrano[4,3,2-de]isoquinolin-3-one.with IC50 = 0.20 µM. heated

VL: PAC (Pharmacological activity); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES 220938-20-5P

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(CA INDEX (preparation of benzpyranoisoguinolinones and related compds. as PARP inhibitors)
220938-20-5 CAPLUS
2H-[1]Benzopyrano[4,3,2-cd]isoindol-2-one, 1,10b-dihydro- (9CI) (CA INDE

Z 2

THERE ARE 567 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT 567

REFERENCE COUNT:

S COPYRIGHT 2007 ACS on STN 2001:772134 CAPLUS 135:318418 CAPLUS L6 ANSWER 3 OF 4 ACCESSION NUMBER: DOCUMENT NUMBER: TITLE:

Guilford Pharmaceuticals Inc., USA U.S., 24 pp., Cont.-in-part of U.S. Ser. No. 922,548. CODEN: USXXAM Preparation of [1]1,10b-dihydrobenzopyrano[4,3,2-delisoindolin-1-one and its analogs as novel poly(ADP-ribose) polymerase (PARP) inhibitors Li, Jia-Her Zhang, Jier Jackson, Paul F.; Maclin, Keith M. PATENT ASSIGNEE(S) SOURCE: INVENTOR (S):

DOCUMENT TYPE: LANGUAGE: FAMILY ACC. NUM. COUNT: PATENT INFORMATION:

PAT	PATENT NO.			KIND		DATE		•	APPLICATION NO.	ICAT	NOI	NO.		ď	DATE	
ns	JS 6306889			E		20011023	023	_	US 1	-866	1998-47502	~		i	9980325	325
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AU 9892982 BR 9812185	EP 1019409	R: AT, BE, (IE, FI	~		JP 2002510332	NZ 503043	NO 2000001001	PRIORITY APPLN. INFO.:				OTHER SOURCE(S): GI

The title compds. [I; Y = alkylhalo, a direct bond, CO, etc.; Z = O, S, NRLI, RI = M, halo, alkyl, etc.); useful for the treatment or prevention of neural or cardiovascular tissue damage related to cerebral ischemia and reperfusion injury in an animal, were prepared Thus, hydrogenating a mixture of Me 9-oxoxanthene-1-actboxylate (preparation given) with NH40Ac and placial AcoH over 10% Pd/C in a bomb at 2000 psi afforded 30% I [Y = a direct bond; X = O; Z = O; Rl-R7 = H]. The 220938-20-5p 9 LL

THE BAC (Biological activity or effector, except adverse); BSU (Biological study, unclassified); SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation); USES (Uses) (Uses); CAPLOS (Uses) (Preparation of [1]1,100-dihydrobenzopyrano[4,3,2-de]isoindolin-1-one and its analogs as novel poly(ADP-ribose) polymerase (PARP) inhibitors) 220338-20-5 CAPLUS (SPE) (CAPLOS (CAPLOS (USES) (CAPLOS (USES) (US

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THERE ARE 345 CITED REFERENCES AVAILABLE FOR THIS RECORD. ALL CITATIONS AVAILABLE IN THE REFORMAT 345 REFERENCE COUNT:

S COPYRIGHT 2007 ACS on STN 1999:184256 CAPLUS L6 ANSWER 4 OF 4 CAPLUS ACCESSION NUMBER: 19

for in,		MKG, KKG, KKG, KKG, KKG, KKG, KKG, KKG,
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130:209714 Tetracyclic heteroaromatic compounds as Tetracyclic heteroaromatic compounds as Listrating neural or cardiovascular tissue damag treating neural or cardiovascular tissue damag treating neural or cardiovascular tissue Gamag Kaith M. Gailford Pharmaceuticals Inc., USA Guilford Pharmaceuticals Inc., USA GODEN: PIXXD2 Parent English	Ň	1189 1189 11,00 11,0
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130:209714 Tetracyclic heteroaromatic poly(ADD-ribose) polymerase treating neutal or cardiova Li, Jia-He; Zhang, Jie; Jac Kaith M. Gollford Pharmaceuticals Ir GOT Int. Appl., 122 pp. CODEN: PIXXD2 Patent English	APPI	MO IN
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130:209714 Tetracycli poly(ADP-r treating n Li, Jia-He Keith M. Guilford P Guilford P CODEN: PIX Parent English	KIND	AU, AZ, LC, LK, LC, LK, LC, LK, LC, LK, LK, LK, LK, LK, LK, LK, LK, LK, LK
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T NU DR(S) ASSI TT TY E: ACC.	PATENT NO	RW RW RW RW 634 CA 2291 CA 2291 CA 2291 CA 229 CA 229 CA 229 CA 229 CA 229 CA 220 CA 2
DOCUMENT NUMBER: TITLE: INVENTOR (S): PATENT ASSIGNEE (SOURCE: DOCUMENT TYPE: LANGUAGE: FARILY ACC. NUM. PATENT INFORMATIC	6	M. AL, M. AL, M. AL, M. AL, M. AL, NO, NO, NO, NO, S10,619 CA 2294133 CA 2294133 CA 2294133 AU 999292 BR 9812185 EP 1019409 EP 1019409 R. AT, HU 20000316, JP 20025103 NZ 50300010 PRICRITY APPLN.
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Title compds. I [Y = alkylhalo, alkyl-COG, COG, direct bond, CO, O, NRII, RS, G = NRIIRE, O, S, CRIZRII, X = NRE, M = NRE, O, S, CRIZRII, S = NRE, M = NRE, O, S = CRIZRII, S = NRE, NRE, M = NRE, O, S = CRIZRII, S = NRE, NRE, M = NRE, OR, CRIZRII, S = NRE, NRE, CRIZRII, S = NRE, NRE, S = NRE, OR, CRIZRII, S = NRE, ΑB

alkyl, C2-C9 alkenyl group, C3-C8 cycloalkyl, C5-C7 cycloalkenyl, aryl, NH2, alkylamino, C02H, or aralkyl) were prepared for use as PARP inhibitors for treating neural or cardiovascular tissue damage. Thus, I [X, Z = 0, Y = NH, R1-R7 = H, the dotted bond is a single bond) was prepared from 9-xanthenecarboxamide by reduction to the amine, conversion to isocyanate, and cyclization and had a PARP-inhibiting IC50 of 0.20 µM.

20938-20-5P

RL: SPN (Synthetic preparation); THU (Therapeutic use); BIOL (Biological study); PREP (Preparation) (USES)
(Preparation of benzopyzanisoquinolinones and benzopyraniophthalazinones as poly(ADP-ribose) polymerase inhibitors)

220938-20-5 CAPLUS

220938-20-5 CAPLUS

220938-20-5 CAPLUS

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REFERENCE COUNT:

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